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Detergent compositions.

 Detergent compositions comprise a special type of oxidation resistant nonionic surfactant and an oxidizing agent which may either be a hypochlorite or a peroxygen material. The surfactant component structurally comprises a C8-C12 alkyl substituted phenoxy hydrophobe alkoxylated with ethylene oxide and/or propylene oxide, with the proviso that the ratio of ethylene oxide to propylene oxide is at least 1 but no higher than 2. Methyl or chloroethyl groups are used to end-cap the surfactant.

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DETERGENT COMPOSITIONS

The present relates to detergent compositions containing surfactants in combination with oxidizing agents.

THE PRIOR ART

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Certain types of cleaning compositions, such as automatic dishwashing detergents, demand the presence of oxidizing agents to operate effectively. Hypochlorite generating compounds are most commonly employed as the oxidizing agent. Although less economical, peroxygen compounds such as sodium perborate have also been reported as useful.

While the present invention has application beyond automatic dishwashing detergent compositions, it is these types of compositions which are of particular commercial concern. Therefore it is appropriate that the problems associated with automatic dishwashing be here set forth.

Automatic dishwashing detergent compositions employ alkaline salts such as sodium silicate, sodium carbonate and sodium tripolyphosphate as the main cleaning agents. A hypochlorite source is generally included in the formulation, mainly for the purpose of breaking up protein soil. Once solubilized, protein soil, derived from foods such as eggs and milk products, gives rise to foaming problems. Foam generation, in turn, interferes with the cleaning action of the machine dishwasher. Without effective foam suppression, the mechanical cleaning action of the dishwasher is reduced because foam build-up partially insulates tableware from the full force of the aqueous washing composition.

Small amounts of nonionic surfactants are usually added to dishwashing compositions to combat the foam problem. Glassware appearance is, to an extent, also improved by the presence of the surfactant. Indeed, spotting and filming is a particularly sensitive criterion by which consumers judge product performance. There is a need for improved performance on glassware.

Numerous types of nonionic surfactants useful in automatic dishwasher detergent compositions have been reported in the literature. For example, U.S. Patent 3,956,401 (Scardera et al.) reports a C7-C10 alcohol alkoxylated to form a three block grouping of oxypropylene/oxyethylene/oxypropylene. U.S. Patent 4,410,447 (Decker et al.) reports a low foaming surfactant using a C7-C11 primary alcohol as a hydrophobe onto which are first attached oxypropylene units followed by a random oxyethylene/oxypropylene mixture. Not only alkyl but also aromatic hydrophobes have been reported. U.S. Patent 4,436,642 (Scott) discloses use of a C6-C12 alkyl substituted phenol alkoxylated first with a block of propylene oxide and then ethylene oxide. Another structural variation has been the incorporation of an end-capping unit to the alkoxylated chain. European Patent Application 0 197 434 (Pruhs et al.) describes defoaming nonionic surfactants formed from the ethoxylation of C8-C18 alcohol end-capped with C1-C4 alkanol, particularly n-butanol.

Most of the aforementioned defoaming surfactants have been formulated for use in powdered automatic dishwashing detergent formulations. More recently, there has been significant consumer interest in pourable liquid versions. Greater challenges are presented by liquid formulations. With powders, many techniques are available to separate surfactant from the oxidizing agent. Providing a coating over the oxidizing materials is, for instance, one method of protecting surfactant against degradation. Even without special separation techniques, powders by their very nature diminish the interaction between components. On the other hand, liquid formulations require their constituents, including any defoaming surfactants to be more oxidatively stable than would ordinarily be necessary in a powder.

SUMMARY OF THE INVENTION

Detergent compositions are disclosed comprising:

(I) From 0.1 to 15 wt.% of a nonionic surfactant identified by formula I

wherein R is an alkyl group containing from 8 to 12 carbon atoms;

EO and PO represent oxyethylene and oxypropylene groups, respectively;

a, b and c may each rang from 0 to 10, with the sum of a, b and c ranging from 6 to 12;

the ratio of EO to PO is at least 1 but no high r than 2; and

Z is selected from methyl or chloroethyl groups and said group is attached to an oxyethylene unit at an oxygen atom thereof;

(ii) from 0.5 to 10 wt.% of an oxidizing agent selected fr m hyp chl rit and hydrogen peroxide generating compounds; and

(iii) from 0 to 80 wt.% of a builder.

The invention also r lates to a m thod of reducing foaming in the cleaning of dishes in an automatic dishwasher comprising contacting the dishes with a bleaching detergent composition containing a nonionic surfactant of formula I.

DETAILED DESCRIPTION OF THE INVENTION

A study of commercially available defoaming surfactants has revealed that certain types of structural features promote oxidative stability and improve cleaning performance. Based on this study, it was determined that the optimum alkoxylate type nonionic surfactant is one having the formula:

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R is an alkyl group containing from 8 to 12 carbon atoms, optimally between 8 and 9 carbon atoms. EO and PO stand for oxyethylene and oxypropylene groups, respectively; EO/PO stands for a random mixture of oxyethylene and oxypropylene units which may range in a ratio from at least 1 but no higher than 2. As represented in the above formula, the notation (EO) and (PO) refer to block polymer units; within the context of the formula the (EO) block may precede or follow the (PO) block depending on the particular surfactant species. Subscripts a, b and c each have a value ranging from 0 to 10, preferably from about 3 to about 10. The sum of a, b and c must be at least 2 and can range up to about 10, optimally from about 6 to about 10. Most importantly, the overall ratio of EO to PO must be preferably between 1 and 2, optimally about 1.5.

End-capped unit Z may either be a methyl or chloroethyl group and these groups are attached to an oxyethylene unit at an oxygen atom.

Surfactants which are particularly preferred are those having the structures II and III outlined below:

$$C_9$$
- C_{12}

$$O(PO)_b(EO) = CH_2CH_2OCH_2CH_2C1$$
(III)

The surfactants used in the present invention may be prepared by condensing an alkyl phenol with propylene oxide and/or ethylene oxide in an amount and respective order dependent upon the particular arrangement of block and random units necessary to form the compound(s). Alkoxylation usually requires the presence of a catalyst which may be sodium or potassium hydroxide, sodium acetate, or preferably an alkali metal alkoxylate such as sodium methoxide. Any other type of catalyst commonly used for alkylene oxide addition reactions with reactive hydrogen compounds may also be employed. These reactions are preferably conducted at elevated temperatures. Upon completion of the alkoxylation, the catalyst may be removed from the reaction mixture by neutralization, filtration or ion exchange.

Methyl groups can be introduced as the end-cap through a method involving reaction between chloromethane and an oxyethylene end unit of a surfactant under conditions of elevated temperature and catalysis. Chloroethyl end-cap groups may be introduced by reaction of an oxyethylen end unit with thionyl chlorid .

Surfactants of the present invention should d sirably have a cloud point below 40°C, preferably less than 20°C, optimally less than about 15°C. Cloud point is defined as the temperature at which clarity of a liquid composition is lost as the xternal temperature is lowered. Lower cloud points are indicative of improved defoaming properties.

Although the surfactants can be used in a wide variety of cleaning products, they are of particular use in automatic dishwasher detergents. Within the autodish category, these surfactants xhibit properties rendering them uniquely suited for the aqueous thixotropic (liquid) form of automatic dishwasher product. The general formulation parameters are set forth in the Table below.

Table A

Automatic Dishwasher Detergent Formulations

10	Component	Powder (wt.%) General Range	Liquid (wt.%) General Range
	Nonionic Surfactant	0.1-10	0.1-10
15	Builder	5-80	5-60
	Sodium Silicate	1-20	1-20
	Filler	0-60	-
20	Bleaching Agent	0.1-20	0.1-20
	Thixotropic Thickener	,	0.5-15
	Water	to 100	to 100

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BUILDER MATERIALS

The dishwashing detergent compositions of this invention can contain all manner of builders commonly taught for use in automatic dishwashing compositions. The builders can include any of the conventional inorganic and organic water-soluble builder salts.

Typical of the well known inorganic builders are the sodium and potassium salts of the following: pyrophosphate, tripolyphosphate, orthophosphate, carbonate, bicarbonate, sesquicarbonate and borate.

Particularly preferred builders can be selected from the group consisting of sodium tripolyphosphate, sodium carbonate, sodium bicarbonate and mixtures thereof. When present in these compositions, sodium tripolyphosphate concentrations will range from about 10% to about 40%, preferably from about 15% to about 40%. Sodium carbonate and bicarbonate when present can range from about 10% to about 50%; preferably from about 20% to about 40%.

Organic detergent builders can also be used in the present invention. They are generally sodium and potassium salts of the following: citrate, nitrilotriacetates, polyacrylates, polyphosphonates, oxydisuccinates, oxydiacetates, carboxymethyloxy succinates, tetracarboxylates, starch and oxidized heteropolymeric polysaccharides. When present, organic builders are preferably present from about 1% to about 35% of the total weight of the detergent composition.

The foregoing detergent builders are meant to illustrate but not limit the types of builder that can be employed in the present invention.

SILICATE

The dishwashing detergent compositions of this invention may contain sodium or potassium silicate. This material is employed as a cleaning ingredient, source of alkalinity, metal corrosion inhibitor and protector of glaze on china tableware. Especially effective is sodium silicate having a ratio of SIO₂:Na₂O of from about 1.0 to about 3.3, preferably from about 2 to about 3.2. Some of the silicate may be in solid form.

OXIDIZING AGENT

A wide variety of oxidizing agents may be employed for use with the dishwashing compositions. Both halogen and peroxygen type materials are encompassed by this invention.

When formulating a liquid automatic dishwashing composition, it is most preferred to employ aqueous sodium hypochlorite as the oxidizing agent. Powder formulations employ halogen donor oxidizing agents in the form of precursor compounds that generate hypochlorite upon addition of water.

Among the suitable halogen donor oxidizing agents are heterocyclic N-bromo and N-chloro imides such as trichlorocyanuric, tribromocyanuric, dibromo- and dichlorocyanuric acids, and salts thereof with water solubilizing cations such as potassium and sodium. An xample of the hydrated dichlorocyanuric acid is Clearon CDB 56, a product manufactured by the Olin Corporation. These oxidants may be employed in admixtures comprising two or more distinct chlorine donors. An xample of a commercial mixed system is a navailable from the Monsanto Chemical Company under the trademark designation "ACL-66" (ACL signifying "available chlorine" and the numerical designation "66", indicating the parts per per und of available chlorine) which comprises a mixture of potassium dichloroisocyanurat (4 parts) and trichloroisocyanurate acid (1 part).

Other N-bromo and N-chloro imides may also be used such as N-brominated and N-chl rinated succinimide, malonimide, phthalimide and naphthalimide. Other compounds include the hydantoins, such as 1,3-dibromo and 1-3-dichloro-5,5-dimethylhydantoin; N-monochloro-C,C-dimethylhydantoin; methylenebis(N-bromo-C,C-dimethylhydantoin); 1,3-dibromo and 1,3-dichloro 5-isobutylhydantoin; 1,3-bromo and 1,3-dichloro 5-methyl-5-ethylhydantoin; 1,3-dibromo and 1,3-dichloro, 5,5-isobutylhydantoin; 1,3-dibromo and 1,3-dichloro 5-methyl-5-n-amylhydantoin, and the like. Further useful hypohalite liberating agents comprise tribromomelamine and trichloromelamine. Dry, particulate, water-soluble anhydrous inorganic salts are likewise suitable for use herein such as lithium, sodium or calcium hypochlorite and hypobromite.

The hypohalite liberating oxidizing agent, may, if desired, be provided in a form of a stable solid complex or hydrate. Examples include sodium p-toluene-sulfo-bromoaminetrihydrate, sodium benzene-sulfo-chloroaminedihydrate, calcium hypobromite tetrahydrate, calcium hypochlorite tetrahydrate, etc. Brominated and chlorinated trisodium phosphate formed by the reaction of the corresponding sodium hypohalite solution with trisodium phosphate (and water if necessary) likewise comprise efficacious materials.

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Preferred chlorinating agents include potassium and sodium dichloroisocyanurate dihydrate, chlorinated trisodium phosphate and calcium hypochlorite. Preferred concentrations of all of these materials should be such that they provide about 0.2 to about 1.5% available chlorine.

Suitable chloride-releasing agents are also disclosed in the ACS monograph entitled "Chlorine-Its Manufacture, Properties and Uses" by Sconce, published by Reinhold in 1962. This book is incorporated by

Among the suitable peroxygen type oxidizing agents are the salts of persulfate, dipersulfate, percarbonate reference. and perborate. Especially preferred are sodium perborate tetrahydrate and sodium perborate monohydrate. Organic peroxy acids such as peracetic acid or 1,12-diperoxydodecanedioic acid may also be employed. Organic peracids are, however, less preferred because of their greater cost.

FILLER

An inert particulate filler material which is water-soluble may also be present. This material should not precipitate calcium or magnesium ions at the filler use level. Suitable for this purpose are organic or inorganic compounds. Organic fillers include sucrose, sucrose esters and urea. Representative inorganic fillers include sodium sulfate, sodium chloride and potassium chloride. A preferred filler is sodium sulfate. Its concentration may range from 0% to 60%, preferably about 10% to 20%.

Minor amounts of various other adjuvants may be present in the detergent powder. These include perfumes, flow control agents, foam depressants, soil suspending agents, antiredeposition agents, anti-tarnish agents, enzymes and other functional additives.

THIXOTROPIC THICKENERS

Thickeners or suspending agents must be added to the liquid versions of automatic dishwasher detergent compositions. They provide thixotropic properties to an aqueous medium. These thickeners may be organic or inorganic water-soluble, water-dispersible or colloid-forming, monomeric or polymeric, and should of course be stable to highly alkaline and oxidative environments. Those especially preferred generally comprise the inorganic, colloid-forming clays of smectite and/or attapulgite types. Smectite clays include montmorillonite (bentonite), hectorite, saponite and laponite clays. Materials of this type are available under trade names such as Thixogel No. 1 and Gelwhite GP from Georgia Kaolin Company (both being montmorillonite). Attapulgite clays include the materials commercially available under the trademark Attagel, i.e. Attagel 40, Attagel 50 and Attagel 150 from Englehardt Minerals and Chemicals Corporation. Mixtures of smectite and attapulgite clays are useful when combined in the weight ratios of 4:1 to 1:5.

Useful thickeners among the organic polymers are water-soluble polycarboxylic acids or salts. Particularly useful is sodium polyacrylate with molecular weight in the range of 1,000 to 50,000, commercially available under the trademark Acrysol and described in GB 2 164 350A (Lai et al.). Preferred amounts of the water-soluble polymeric carboxylic acid will range from about 0.01 to about 3%.

Amounts of water present in the liquid type compositions should neither be so high as to produce unduly low viscosity and fluidity, nor so low as to produce unduly high viscosity and low flowability, thixotropic properties in either case being diminished or destroyed. Water will generally be present in an amount ranging from 45 to 75 wt.%, preferably about 55 to 65 wt.%.

The following examples will more fully illustrate the embodiments of this invention. All parts, percentages and proportions referred to herein and in the appended claims are by weight unless otherwise indicated.

Interactions of nonionic surfactants with hypochlorite are complicated. Attack sites are the ether linkages, terminal hydroxyl group and the hydrophobic alcohol unit. Each of these structural components of typical commercially available nonionic surfactants has been investigated.

A liquid type automatic dishwashing detergent base formulation is outlined in Table I. Various commercially known defoaming surfactants (1-16) and an experimental surfactant (17), listed in Table II, were incorporated at 2 wt.% to the base liquid composition. Samples were stored in glass vials and submerged in a temperature controlled water bath. Subsequ at thereto, the samples were titrated for available chi rine and the pH analyzed.

Table I

Liquid type Automatic Dishwashing Detergent Base

5	Component	Weight %
	Sodium	16.0
	Tripolyphosphate	
	Sodium Carbonate	6.0
10	Sodium Silicate	8.0
	Sodium Hydroxide	1.2
	% Available Chlorine	1.0
	(as hypochlorite)	
	Bentonite clay	4.0
15	Attapulgite clay	1.0
	Water	Balance

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Table II

Surfactants Investigated

	Sur-	Chemical Structure
25	fac- tant	
30	2 [(3 C 4 C 5 C	kanol-(PO)4(EO)4(PO)4 alkyl)(SO3Na)C6H3]-O-]C6H4(SO3Na)] sH5O-(EO)16.5(PO)11 sH5O-(EO)4.5(PO)12 12-C15 alkanol-(EO)7
<i>35</i>	7 C 8 C 9 C	3 alkanol-(EO)5-CH2COO-Na ⁺ 5H5O-(EO)2-CH2COO-Na ⁺ 5H5O-(EO)3-CH2COO-Na ⁺ 5H5O-(EO)4-CH2COO-Na ⁺
40	11 (0 12 C 13 C 14 C	BH5O-(EO)6-CH2COO-Na ⁺ C ₈ alkyl)C ₆ H4O-(EO)4-CH2COO-Na ⁺ 12-C ₁₄ alkanol-(EO)6 12-C ₁₄ alkanol-(EO)6-CH3 16-C ₁₈ alkanol-(EO)45
45	16 C	₁₆ -C ₁₈ alkanol-(EO)4.5-CH3 ₁₂ -C ₁₅ alkanol-(EO)8-C(CH3)3 ₆ -C ₁₀ alkanol-(PO)4(EO)4(PO)4CH2C6H5

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Table ill Percent Available Chlorine

					
Surfactant	1 v	veek	<u>4 we</u>		5
Odridota	25°C	37°C	<u>25° C</u>	<u>37°C</u>	5
Base alone	1.00	0.95	1.00	0.87	
1	0.60	0.35	0.29	0.035	
2	0.95	0.52	0.90	0.67	
3	0.65	0.39	0.45	0.14	10
4	0.81	0.53	0.60	0.25	
5	0.54	0.17	0.14	0.00	
6	0.91	0.69	0.64	0.19	
7	0.74	0.54	0.61	0.25	15
8	0.78	0.48	0.66	0.32	
9	0.98	0.44	0.80	0.50	
10	0.93	0.46	0.79	0.48	
11 .	0.92	0.50	0.81	0.50	
12	0.87	0.45	0.50	0.15	20
13	0.94	0.80	0.61	0.20	
14	0.87	0.80	0.60	0.15	
15	0.86	0.85	0.65	0.25	
16	0.85	0.24	0.25	0.00	25
17	0.75	0.50	0.40	0.10	25

Certain conclusions can be drawn from the chlorine stability results outlined in Table III. After storage for a month, the surfactant which least interacts with chlorine, and thereby allows for a higher available chlorine content, is surfactant 2. Indeed, the surfactant is widely used commercially for defoaming autodish compositions; it is sold by DOW under the trademark Dowfax 3B-2. The aromatic structures at either end of the ether linkage undoubtedly contribute to this stability. Surfactants 9, 10 and 11 exhibited the next best stability. Performance of these materials compared favourably with that of surfactants 3 and 4 which are not end-capped with an acetate group. Comparison of stability between surfactants 1 and 17 further illustrates the effect of end-capping. Surfactant 17 is substantially identical to surfactant 1 with the exception of a benzyl end-group. From these results, it would appear that end-capping is highly beneficial for protection against oxidative degradation.

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Another observation is the better performance of surfactant 4 than that of surfactant 1. Although the structures vary in several respects, it would appear that the greatest influence is that derived by the hydrophobe portion. Surfactant 1 is based on a C6-C10 alkanol while surfactant 4 is based on phenol. The phenolic hydrophobe has better stability and interferes less with the available chlorine.

Based on the foregoing results, the preferred defoaming surfactant should be a molecule with an aromatic hydrophobe, and protected at this terminal hydroxyl group with an end-capping unit.

EXAMPLE 2

Not only must surfactants for automatic dishwasher compositions be chlorine stable, they must, most importantly, deliver defoaming and wetting action. Foam measurements on many of the surfactants listed in Table II were performed to 500 ml aqueous solutions containing 0.6 wt.% surfactant. The foam testing device consisted of a Waring blender surrounded by a jacketed column to maintain temperature. Foam heights were measured after 60 seconds of agitation and after 60 seconds at rest.

Table IV Surfactant Foaming Assessment

		
Surfactant	Initial Foam Height (mm)	Final Foam Height (mm)
1	trace	2.5
2	54	48
3	5	15
4	trace	trace
5	55	48
6	7	10

Th data show that surfactant 2, with its highly stable structure, unfortunately is relatively poor at defoaming. A comparis n of surfactants 3 and 4 indicates that there is a significant defoaming benefit where the amount of ethylene oxide is minimized and the presence of propylene oxide maximized. A confirmatory result in se n when surfactants 1 and 5 are compared, the former having an excess of propylene oxide and the latter containing only ethylene oxide. Surfactant 1 had substantially better defoaming performance.

In Examples 3 and 4, the advantages of an excess of propylene oxide are mitigated by including end-cap

units necessary for increased bleach stability.

EXAMPLE 3

A number of further surfactants were evaluated to determine the optimum structural properties required for use in automatic dishwasher compositions containing chlorine. Polytergent SLF-18, a product of the Olin Corporation, was used as a reference. This material is known to be a C7-C10 alcohol alkoxylated to form a three block grouping of PO/EO/PO ending in a hydroxyl group. Test surfactants were based on structure IV outlined

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$$(CH_3)_3CC_6H_4O-(PO)_b(EO)_a$$

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IV

Table V

25	Sample No.	<u>b,a</u>	<u>z</u>	Hydroxyl No.	Molecular Weight	HLB
	Surfactant					
	18	4.	4 H	106.0	559	5.05
	19	·	5 H	93.0	647	5.75
30	20		4 CH ₃	7.5	573	3.95
	21	4.		8.0	650	1.10
			· · · · · · · · · · · · · · · · · · ·	2.0	738	1.80
	22	4,	0 CH2C6H5	2.0	1800	4.55
	SLF-18	•			1866	4.00

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The surfactants of Table V were evaluated for a number of physical properties. Cloud point of surfactants in water and in an electrolyte solution are reported in Table VI. Cloud point values were determined by preparing solutions of 0.1 grams surfactant in 100 ml distilled water and a similar concentration in an electrolyte solution. The latter was formulated to simulate levels and types of builder salts in a typical wash liquor. The electrolyte combination of materials were used at a strength of 4 grams per 1,000 ml water at pH 10.5 and included sodium tripolyphosphate/sodium carbonate/sodium polysilicate at a ratio of 55/33/12.

Table VI

45	Cloud Points (°C)						
	Surfactant	Distilled Water	Electrolyte Solution				
	18	18	<5				
50	19	45	· < 5				
	20	<5	<5				
	21	<5	<5				
	22	<5	<5				
55	SLF-18	19	<5				

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As the temperature of a nonionic solution is raised the reduced hydration of the ethylene oxide groups results in the formation of two separate isotropic phases. Onset of this phenomenon is described as the cloud point. For phase theoretical considerations, this is a liquid/liquid separation with a lower critical solution temperature.

The extent of this phas behaviour is a critical function of the foam volume generated or suppressed by the nonionic. Foam volumes fall drastically above the cloud point and create conditions where the capacity to limit food soil foams is greatly enhanced. Inasmuch as the wetting ability of the surfactant is a function of the concentration in the aqueous phase it is important that the cloud point not be so I was to limit this function. Consequently, there is a need to balance both of these conditions by selecting surfactants with appropriate cloud points given the operating temperatures found in most dishwashing machines.

From Tabl VI, it can be seen that cloud points, as measured in distilled water, are quite satisfactory for surfactants 20-22 which are end-capped with methyl and benzyl groups. On the other hand, surfactants that are not end-capped such as 18, 19 and SLF-18, provided unacceptable cloud points.

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Table VII reports foam height measur ments made under machine wash conditions with and without the presence of soil. The test procedure was similar to that reported in Example 2. Here however, ther was also added 2.0 grams of an automatic dishwashing liquid base formulation for alkalinity purposes along with 0.02 grams of a surfactant in 500 ml tap water, which combination represents home dishwasher conditions of 40.0 grams detergent per 10 liter wash. Foam heights were measured in millimeters after one minute of agitation followed by one minute of quiescence.

Table VII

Foaming As	ssessment Under Automatic W	ashing/
	Machine Conditions	

·	Machine Conditions	•	•		
Surfactant	Conditions				
	No Soil	Soil			
18	9	11			-
19	10	11			20
20	5	10	•		
21	2	9			
22	2	9		•	
SLF-18	0	5			25
No surfactant	-	7			

Table VIII reports results of surface tension measurements on six surfactants in electrolyte solution. Using a Cahn electrobalance and a Wihelmy plate setup, values of surface tension as a function of concentration were measured at 45°C. Isotherms resulting therefrom were plotted as surface pressure versus log molarity. Relevant physical data were derived from these curves.

Table VIII

Sur	face Tension	Measureme	ent
Surfactant	CMC	<u>cmc</u>	<u>σ</u> ,
18	398	38.00	90.6
19	562	38.25	99.1
20	31.6	36.50	101.1
21	44.7	29.75	135.9
22	39.8	30.25	140.4
SLF-18	1.00	32.25	69.0
PARAMETE	R UNITS		
CMC ×10	-6M		
πcmc erg	s/cm²		
σ Ų/mole	ecule		

At very low concentration, surface pressure is negligible and increases very slowly. As concentration builds, a point is reached where pressure increases dramatically up to an inflection point known as the critical micelle concentration (CMC). Very little increase in pressure occurs thereafter. This CMC represents a range of concentration where continued adsorption of surfactants at the interface is minimal and the formation of aggregates of surfactants in the bulk liquid initiates micellization. Surfactants that generate high surface pressures are regarded as being highly effective. Regardless of the amount needed, it is the absolute value of surface tension reduction that quantifies the effectiveness of a surfactant. However, that amount needed to achieve maximum surface pressure is the CMC and it is a measure of efficiency.

With these terms defined above, it will be seen that SLF-18, the control material, is much more efficient, its CMC is at a low r concentrati n, than benzyl-capped derivativ s.

Table IX reports hypochl rite stability values. In these evaluations, each surfactant is dispersed in a base formula of a typical automatic dishwashing liquid so that there are equimolar solutions equivalent to 2 weight % of SLF-18. Initial available chlorine level was adjusted to 1.0%. Each week, samples wer taken and titrated for available chlorine including a surfactant-free case and one with SLF-18.

Table IX
Percent Available Chlorine

5	Week No.			25°C S	urfactant		
_		<u>18</u>	<u>19</u>	<u>20</u>	22	SLF-18	None
	0	1.000	1.000	1.000	1.000	1.000	1.000
	1	0.784	0.818	0.900	0.753	0.256	0.986
10	. 2	0.661		0.884	0.683	0.154	0.971
	4	0.578	0.554	0.771	0.572	0.063	0.941
	6	0.456	0.421	0.628	0.478	0.024	0.931
	8	0.402	0.367	0.541	0.417	0.007	0.889
	Week No.	45°C Surfactant					
15		18	<u>19</u>	20	22	SLF-18	None
	.0	1.000	1.000	1.000	1.000	1.000	1.000
	, 1	0.480	0.425	0.667	0.535	0.027	0.865
20	2	0.277	0.237	0.466	0.275	0.000	0.767
	4	0.060	0.049	0.151	0.083	0.000	0.635
	6	0.050	0.037	0.113	0.057	0.000	0.468
	8	0.002	0.019	0.043	0.017	0.000	0.387

Hypochlorite stability test results of Table IX show that the methyl end-cap material (20) has the greatest stability.

Use of a benzylic end-cap does not result in optimum surface properties. Without end-capping, as seen with SLF-18, stability is severely sacrificed since the ether linkages are now subject to oxidation.

30 EXAMPLE 4

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The experiments that follow detail work on surfactants similar to that reported on in Example 3 but now replacing the tert-butyl with a nonyl group on the phenol portion.

Table C outlines the nonyl phenol derivatives whose structure V is set forth below:

nony1—
$$C_{6}H_{4}O$$
—(PO)_b (EO) $\frac{}{a}$

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Table X

V

45	Sample No. Surfactant	<u>b,a</u>	<u>z</u>	Molecular Weight
	23	4,4	Н	628
50	24	4,6	H	716
	25	4,8	Н	804
	26	0,6	Н	484
	27	4,4	CH ₃	642
	28	4,6	CH ₃	730
<i>55</i>	29	4,8	CH ₃	818
	30	0,6	CH₃	498
	SLF-18	-	Н	1800

The surfactant of Table X were evaluated for a number of physical properties. Cloud points of these materials are list d in Table XI.

Table XI

Cloud Points (°C)

	<u> </u>	
Surfactant	Distilled Water	Elec
23	<0	
24	28	
25	43	3
26	<0	<0
27	3 5	30
28	37	31
29	46	40
30	<0	<0
SLF-18	19	<0

Table XII reports foam height measurements made under machine wash conditions with and without the presence of soil.

The test procedure was similar to that reported in Example 3. Although the foam measurement is more qualitative than quantative, it is useful in discriminating among various materials. In this case, SLF-18 appears best in foam suppression. However, all of the samples in the Table are much better than typical anionic defoamers such as Dowfax 2A1 with foam heights of 25 to 30.

Table XII

Foaming Assessment Under Automatic Washing

Machine Conditions

•		_
Surfactant	Condi	itions
	No Soil	Soil
23	3.5	8.5
24	4.0	9.0
25	5.5	9.0
26	3.5	9.5
27	3.0	8.0
28	4.0	8.5
29	6.5	8.0
30	4.5	7.5
SLF-18	0.0	4.0
No surfactant	0.0	7.0

Table XIII reports results of surface tension measurements on the nonyl phenol derivatives. Values reported in this Table were obtained by the method already outlined in Example 3.

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Tabl XIII
Surface Tension Measurement

	Surfactant	CMC	πεπε	<u>σ</u>
5	23	1.58	32.25	48.98
	24	1.78	32.00	52.93
	25	3.55	31.50	56.13
	26	14.1	36.75	53.65
10	27	2.11	34.00	34.20
	28	2.51	34.00	47.58
	29	2.82	33.25	55.50 .
	30	18.8	39.50	51.20
15	SLF-18	1.00	32.25	69.0
13	PARAMETER	UNITS		

CMC x10-6M

πcmc ergs/cm²

20 σ Å²/molecule

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A discussion of surface tension measurements and their significance has previously been presented under Example 3 and is not here repeated. From that discussion, it is to be understood that the larger the CMC value, the more efficient is the surfactant. From Table XIII it is evident that several of the sample surfactants of this invention come very close in CMC value to SLF-18. Samples 23-25 and 27-29 all had CMC values very close to that of SLF-18. These were all considerably better than the CMC values of the tert-butyl phenol derivatives listed in Table VIII. Further, it is noted that samples 26 and 30 which were wholly ethoxylated and contained no proproxylation had significantly poorly CMC values. Thus, it is evident that there must be an upper limit to ethoxylation; some propylene oxide must be present within the molecule.

Example 5 demonstrates the improved oxidative stability of the non-ionic surfactants according to the present application.

EXAMPLE 5

The base composition according to Example 1 was used. A variety of different surfactants were dispensed therein and titrated for available chlorine including a control surfactant-free case and one with SLF-18 (commercial surfactant).

nonyi - C₆H₄O - (PO)_b(EO)_a - Z

40	Sample No. Surfactant	<u>b,a</u>		<u>z</u>
	23	4,	4 H	
	24	4,	6 H	
45	· 25	4,	8 H	
	26	0,	6 H	
	27	4,	4 CH:	3
	28	4,	6 CH;	3
	29	4,	8 CH;	3
<i>50</i>	30	0,	6 CH:	3

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Table

			% Available	Chlorine				
Week No.	23	<u>25</u>	<u>27</u> ·	25°C <u>29</u>	30	SLF-18	None	5
. 1	0.82	0.80	0.92	0.87	0.89	0.25	0.98	
2	0.71	0.68	0.85	0.80	0.88	0.15	0.97	
4	0.64	0.58	0.80	0.72	0.83	0.06	0.94	
6	0.57	0.51	0.78	0.62	0.78	0.02	0.93	10
8	0.52	0.44	0.73	0.55 45°C	0.70	0.01	0.88	
1	0.55	0.45	0.78	0.65	0.72	0.027	0.86	
2	0.32	0.23	0.52	0.34	0.52	0.00	0.76	15
4	0.10	0.06	0.33	0.13	0.31	0.00	0.65	
6	0.06	0.00	0.11	0.06	0.19	0.00	0.46	
8	0.00	0.00	0.06	0.00	0.04	0.00	0.38	

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ΔN

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Compositions formulated with surfactants 23-26 can be seen from the Table as retaining less available chlorine, for any given test week, than those of 27-30. Thus, the data demonstrate the effectiveness of end-capping, especially with methyl groups.

Degree of alkoxylation is another important factor as shown by the Table. Thus, advantage is gained where the PO:EO ratio is 1:1 or 1:1.5 compared with that of 1:2. Compare surfactant compositions 23/24 and 27/28 against 25 and 29, respectively.

From the Table it is also evident that any of surfactant compositions 23-30 are an improvement over the common commercially available SLF-18 identified in the specification as a C7-C10 alcohol alkoxylated to form a three block grouping of PO/EO/PO ending in a hydroxyl group.

Surfactant compositions 26 and 30 having only EO type alkoxylation do have similar oxidation stability to that of the mixed PO/EO alkoxylated nonylphenols. However, these two surfactants have exceptionally poor surfactant properties as demonstrated in Table XIII at page 14 of applicants' Preliminary Amendment. From that Table, one can see that surfactants 26 and 30 have a CMC (critical micelle concentration) one order of magnitude inferior to that of surfactants 23-25 and 27-29.

From the foregoing experiments it can be inferred that alkoxylated C₈-C₁₂ alkylphenols have excellent oxidative stability. Especially preferred are structures where alkoxylation is a mixture of EO/PO of at least 1 but no higher than 2, preferably being a ratio of from 1 to 1.5. Additionally, it is advantageous to have a methyl end-capping unit.

Claims

1. A detergent compositions comprising:

(i) from 0.1 to 15 wt.% of a nonionic surfactant identified by formula I:

wherein R is an alkyl group containing from 8 to 12 carbon atoms;

EO and PO represent oxyethylene and oxypropylene groups, respectively;

a, b and c may each range from 0 to 10, with the sum of a, b and c ranging from 6 to 12;

the ratio of EO to PO is at least 1 but no higher than 2; and

Z is selected from methyl or chloroethyl groups and said group is attached to an oxyethylene unit at an oxygen atom thereof;

(ii) from 0.5 to 10 wt.% of an oxidizing agent selected from hypochlorite and hydrogen peroxide generating compounds; and

(iii) from 0 to 80 wt.% of a builder.

2. A composition as claimed in claim 1 wherein R is a C₈ or C₉ alkyl radical.

3. A composition as claimed in claim 1 or claim 2 wherein Z is a m thyl r chloroethyl group.

4. A composition as claimed in any one of claims 1 to 3 wherein the ratio of EO to PO is between 1 and 65

5	 5. A composition as classificate. 6. A composition as creleasing compound selection as classificate. 7. A composition as classificate. 9. Percentage of the composition as	laimed in any or cted from the gra im dichloroisocy laimed in any or apound selected	ne of claims 1 oup consisting of anurate dihydra ne of claims 1 to d from the g	to 6 whereir of sodium hy te, and mixtu o 5 wherein roup consi	n the oxidizing pochlorite, so the state of the oxidizing sting of dipositions.	g agent is a dium dichlo agent is a ersulfate, p	a chlorine proisocya- hydrogen persulfate,
10	8. A composition as classification as classifica	laimed in claim ydrate. imed in any one c imed in any one c imed in claim 10 t	of claims 1 to 8 v of claims 1 to 8 v further comprisi	vhich is in gra vhich is in a l	anular form. iquid form. opic thickener.		
15	12. A composition as cl attapulgite type clays or w 13. A method of washin claimed in any one of claim	laimed in claim ater-soluble pol ng dishes compi	11 wherein said ymeric carboxylising contacting	i thickener i ic acid or sal g the dishe:	s a clay seled t thereof.	ted from s	
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- Detergent compositions.
- Detergent compositions comprise a special type of oxidation resistant nonionic surfactant and an oxidizing agent which may either be a hypochlorite or a peroxygen material. The surfactant component structurally comprises a C₈-C₁₂ alkyl substituted phenoxy hydrophobe alkoxylated with ethylene oxide and/or propylene oxide, with the proviso that the ratio of ethylene oxide to propylene oxide is at least 1 but no higher than 2. Methyl or chloroethyl groups are used to end-cap the surfactant.



European
Patent Office

EUROPEAN SEARCH REPORT **Application Number**

EP 89 30 3614

	OCUMENTS CONS				
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	The present search report has	been drawn up for all claims			
	Place of search	Date of completion of s	earch	<u> </u>	Examiner
	The Hague	17 December 9	1		FISCHER W.H.F.
Y: po de A: te O: no P: In	CATEGORY OF ITED DOCS articularly relevant if taken alone articularly relevant if combined wit ocument of the same catagory achnological background on-written disclosure itermediate document beory or principle underlying the in	h another	the filing of D: document L: document	late cited in the cited for of the same p	her reasons